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Liposome Targeting to Tumors using Vitamin and Growth Factor Receptors

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Liposome-encapsulated anticancer drugs reveal their potential for agents to solid tumors. Advances in liposome technology have redue to their ability to enhance the delivery of chemotherapeutic due to their high expression levels on various forms of cancer and selectively increasing the efficacy of carried agents against receptorsulted in the development of ligand-targeted liposomes capable of increased therapeutic efficacy and decreased nonspecific toxicities become attractive targets for ligand-directed liposomal therapies bearing tumor cells. Receptors for vitamins and growth factors have

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their ability to internalize after binding to the liposomes conjugated to receptors' natural ligands (vitamins) or synthetic agonists (receptor-specific antibodies and synthetic peptides). This chapter summarizes various strategies and advances in targeting liposomes to vitamin and growth factor receptors in vitro and in vivo with special emphasis on two extensively studied liposome-targeting systems utilizing folate receptor and HER2/neu growth factor receptor.

I. INTRODUCTION

Growth, nutrition, and differentiation are among the key functions of living cells that constitute our body. Energy-rich nutrients and structural building blocks, such as sugars, fats, and amino acids, are needed in abundance and enter cells from a lavish extracellular pool. To effectively metabolize the nutrients and build its own bulk, the cell needs vitamins and other enzyme cofactors that are not produced by the cell itself and must be absorbed from a relatively scant environment. Raw power of cellular growth provided by metabolism of nutrients is tamed by the process of cell growth control and differentiation based on intricate communication between the often remote groups of cells through hormones, chemical effectors, and growth factors. Cellular uptake of vitamins and response to hormones and effectors depends on receptor proteins that specifically interact with these substances and elicit proper physiological responses on the cellular level.

Growth and differentiation of the cells are the first functions to change dramatically when the cells become malignant. In malignant cells, the molecular machinery of hormone and growth factor receptors is changed to provide constant stimulation of unabridged cell growth and reduced ability for normal differentiation. Intensive biosynthesis of cellular components, in turn, requires increased supply of metabolic cofactors. Thus, normal and malignant cells often have profound differences in the abundance and function of vitamin and growth factor receptors. Because of these differences and because the receptors for water-soluble, hydrophilic molecules such as peptide hormones and vitamins are usually exposed at the cell surface, vitamin and growth factor receptors are promising "recognition tags" for targeted anticancer drug delivery.

In the malignant phenotype the expression levels of certain vitamin and growth factor receptors are often substantially elevated (Sporn and Roberts, 1985; Slamon et al., 1987, 1989; Berchuck et al., 1990; Weitman et al., 1992; Ross et al., 1994; Fan and Mendelsohn, 1998).

used to target a variety of toxins (Leamon and Low, 1992; Leamon et al., ties to cell-surface receptors for vitamins and growth factors have been Antibodies, antibody fragments, and small molecule ligands with affini Jinno et al., 1996; Lu et al., 1999) and small-molecule therapeutics cally to receptor-overexpressing cancer cells. There are some limitations (Sivam et al., 1995; Uckun et al., 1998; Tolcher et al., 1999) specifi (Hartman et al., 1994; Wilder et al., 1996; Multani et al., 1998; Iznaga-1993; Ramakrishnan et al., 1996; Rosenblum et al., 1999), radionuclides hypoalbuminemia, weight gain, hypotension, and peripheral and pulicity in this case is vascular leak syndrome (VLS), characterized by nonspecific toxicity characteristic for the toxin domain. Frequent toxage quite immunogenically and even at subtherapeutic doses still have inherent to each of these approaches Toxin conjugates, for example Escobar, 1998; Wilbur et al., 1999), enzymes (Rodrigues et al., 1995; systems are limited by their rapid clearance from the circulation, low calculations and an on-site radiopharmacy (Multani and Grossbard, to use in large clinical centers due to the need for complex dosimetry barrier in solid tumors (Weinstein et al., 1987), and may be limited icant hematological toxicities, are limited by the so-called binding-site the treatment of hematological cancers but are associated with signifradionuclide-antibody conjugates have shown significant promise in 1993; Sausville et al., 1995; Stone et al., 1996). Cancer cell-specific the vascular compartment (Vitetta et al., 1991; Grossbard et al., 1992 monary edema resulting from extravasation of fluid and proteins from sure to the biological media after administration into the body. These number of active drug delivered per targeting event (interaction of the the cytotoxic agent upon coupling to a targeting ligand or upon expotargeted drug carrier with the target cell), and potential inactivation of on liposomes. limitations, however, are not inherent to drug-delivery systems based 1998; Iznaga-Escobar, 1998). All these types of targeted drug-delivery

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Liposomes were first described in 1960s when Bangham and co-workers discovered that lecithin swelling in aqueous buffers forms microscopic bodies composed of nested lipid bilayers enclosing aqueous interior (Bangham, 1963; Bangham et al., 1965). During more than 3 decades since this discovery, liposomes have been the subject of numerous studies, books, and reviews. For details of current liposome technology we refer the reader to an excellent and comprehensive textbook by Lasic (1993) and to a recent book that offers state-of-the-art coverage of liposome drug delivery in general (Lasic and Papahadjopoulos, 1998).

Generally, liposomes are vesicular structures consisting of one or more enclosed lipid-bilayer membranes (lamellae) that encircle an

LIPOSOME TARGETING TO TUMORS

aqueous space containing the solute of interest, which in the context of this chapter is an anticancer active principle, whether it is a small-molecular-weight drug or a large macromolecule line DNA. Liposomes are generally produced when certain lipids, especially the natural lipid components of biomembranes, e.g., phosphatidylcholine, are allowed to swell in aqueous buffers and then are fragmented by mechanical shearing, ultrasonication, microfluidization, or extrusion through micro-or nanoporous membranes; alternatively, the lipids are solubilized in the presence of a dialyzable detergent which is then removed to allow the lipid molecules to associate in bilayers and form vesicles. Lipids in the membranes of liposomes are organized in two-dimensionally oriented (liquid crystalline) phases not unlike biological membranes; this property of liposomes prompted their use as a model of biomembranes in many biophysical and biochemical studies (Lasic, 1993).

The microcontainer nature of liposomes and their natural compatibility with living tissues makes them ideally suited as drug carriers. Water-soluble drugs can be loaded into liposomes simply by sequestering (entrapment) drug solution within liposomes during their formation. Moderately lipophilic drugs having the properties of weak acids or bases can be very effectively encapsulated by "active" or "remote" loading methods by creating a transmembrane gradient of pH and/or electrochemical potential that "drives" the drug into the liposome (Nichols and Deamer, 1976; Mayer et al., 1985; Haran et al., 1993). More lipophilic drugs with poor water solubility usually associate with the liposome bilayer.

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spect to size (from 30 nm to several micrometers), number of bilayers commonly formulated into unilamellar liposomes 70-150 nm in size change of membrane components out of the liposomal membrane in encapsulated drug in the circulation, and because it limits the exmulations because it reduces membrane permeability (Papahadjopoucholesterol. Cholesterol is an important component of liposomal forare phospholipids, such as phosphatidylcholine, and sterols, such as Common lipid components of liposome membranes, as mentioned above, tially improve the fate of drug-carrying liposomes in the body (Fig. 1). the surface-attached sugar or hydrophilic polymers which may substan-(unilamellar, oligolamellar, or multilamellar), lipid composition, surbilayer organization of lipids, various types of liposomes differ with rewhich allows them to permeate through the vasculature of tumors but blood plasma (Allen, 1981; Damen et al., 1981). Anticancer drugs are los et al., 1972; Mayhew et al., 1979), and thus increases stability of face charge, and presence or absence of so-called steric stabilization by Despite common structural features such as vesicular structure and

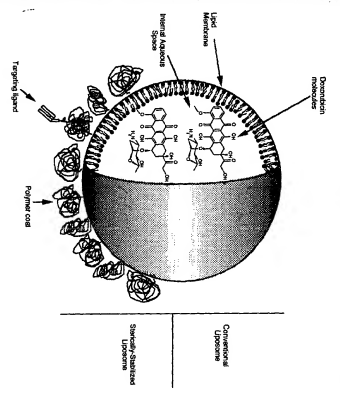


Fig. 1. Schematic representation of a drug-loaded liposome both with poly(ethylene glycol) coating (sterically stabilized liposome, SSL) or without coating ("conventional" liposome). The liposome has a lipid bilayer membrane that encapsulates internal aqueous space used to hold drug substances. Some amphiphilic drugs, such as doxorubicin, can be encapsulated at concentrations exceeding their aqueous solubility and form crystals in the liposome interior. Yet more hydrophobic drugs may be carried within the lipid bilayer. Further modification of the surface by covalently attaching targeting ligands such as small-molecule ligands (e.g., folic aid) or antibody fragments can result in liposomes that are specifically endocytosed by target cells that express a receptor for that ligand or an antigen for the antibody, i.e., folate receptor or growth factor receptor. Phospholipids, such as distearcylphosphatidylcholine (DSPC) and other phosphatidylcholines (lecithins), as well as cholesterol, are common components of systemically delivered liposome formulations, although other lipid compositions are possible.

not of normal tissues (Hobbs et al., 1997). These liposomes are administered intravenously and should be able to persist in the circulation long enough to allow liposome extravasation into the tumor. Early liposomal formulations suffered from fast clearance from circulation by the cells of mononuclear phagocytic system. Significant advance in liposome technology was the discovery that coating of liposome surface with some oligosaccharides or polymers, especially poly(ethylene glycol)

("sterically stabilized liposomes") results in remarkably low blood-clearance rates (Gabizon and Papahadjopoulos, 1988; Klibanov et al., 1990; Allen et al., 1991; Papahadjopoulos et al., 1991, Woodle and Lasic, 1992). This discovery, along with the development of highly efficient methods of "active" loading of drugs into liposomes (Cullis et al., 1997) aided in the development of liposomal drugs all the way to clinic.

associated with the free drug (Muggia et al., 1997; Ranson et al., 1997; when compared to the free drug, while significantly altering the toxcristine have shown either similar or increased therapeutic efficacy of the drug biodistribution is possible because of the drug persistence tumor tissue-specific biodistribution of the drug-loaded liposomes for a result from a variety of factors, the major one being the substitution of et al., 1999; Shapiro et al., 1999; Valero et al., 1999). These benefits Gabizon, 1998; Northfelt et al., 1998; Drummond et al., 1999; Gelmon icity profile and reducing many of the common nonspecific toxicities et al., 1999; Gabizon and Barenholz, 1999). The net effect is an increased et al., 1997; Allen, 1998; Bally et al., 1998; Martin, 1998; Drummond Allen and Papahadjopoulos, 1993; Allen et al., 1995a). Such alteration relatively nonspecific biodistribution of the drug itself (Hwang, 1987; and selective extravasation of drug-loaded liposomes at the tumor site mors which lack lymphatic drainage (for recent reviews see Gabizon vessels for microparticles, and low clearance rate of liposomes from tuwithin the circulating liposomes, higher vascular permeability of tumor and an increased accumulation of liposomal drugs at the tumor. This and benefits further "active," ligand-directed targeting of drug-loaded "passive" targeting of liposomes to solid tumors not only increases the liposomes to cancer cells. therapeutic index of liposome-loaded anticancer drugs, but also enables Clinical trials with liposomal drugs such as doxorubicin and vin-

resides their propensity for extravasation into malignant rather than normal tissues, liposomes offer several other advantages as drug carriers for ligand-directed targeting (Rosenberg et al., 1987; Lee and Low, 1994, 1995; Park et al., 1995, 1997a, 1998a; Kirpotin et al., 1997a, 1998). Liposomes can deliver much larger drug payloads per each targeting event. For example, one doxorubicin-loaded liposome internalized into cancer cells via a ligand-receptor interaction carries into the cell approximately 2×10^5 molecules of the drug. Liposomes can protect the encapsulated drug from degradation by enzymes and neutralization by antibodies in the central compartment until the drug enters the target

Ligand-directed targeting may increase the bioavaliability of liposomal drug to target cells. For the drug to work, it must be released from

the carrier upon reaching the target tissue. Following extravasation, mulations, such as Doxil (Alza Corp.) the drug is released slowly in surrounding the cancer cells. From relatively stable doxorubicin fornontargeted liposomes are primarily found in the tumor interstitium surface. If the liposome is targeted via conjugated ligands to internal diffuse into the nearby cancer cells. Liposomes may also be modified the area of close proximity to the tumor, where the free drug may then lent or noncovalent conjugation of a targeting ligand to the liposome for targeting to specific ligands on various target cells by either covacreasing the amount of bioavailable drug (Machy et al., 1982; Lee et al., tion, liposome internalization may simply serve to limit the diffusion drugs, however, such as cytosine arabinoside, may be degraded, result kaloids, can escape the organelles of endocytic pathway intact; other liposomal delivery, such as doxorubicin, daunorubicin, and vinca al lysosomes and in the presence of degradative enzymes located in these drug must be stable in the acidic environment of the endosomes and liposome and release of the drug occurs intracellularly, effectively inizing receptors located preferentially on cancer cells, breakdown of the of the released drug away from the tumor, thus exposing more of the ing in significantly diminished activity (Huang et al., 1983). In addiintracellular organelles. Luckily, current anticancer drugs of choice for 1998; Drummond et al., 1999). For this approach to be effective, the tumor to the cytostatic agent (Allen et al., 1998). Active targeting of liinternalizing CD19 protein exposed on malignant B-cells in vivo (Lopes targeting approach used in treating solid tumors. Allen and co-workers blood-borne malignant cells will be unable to benefit from the passive posomes will also be important in treating hematological cancers where de Menezes et al., 1998). have recently demonstrated encouraging results targeting liposomes to

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The above considerations indicate that the efficacy of ligand-directed liposome targeting is higher if after binding to the surface of the target cell the liposome becomes internalized by this cell. Because vitamin receptors often perform cellular transport function and growth factor-receptor complexes are often internalized by the cell as part of cellular response it is more likely that liposomes conjugated to ligands specific to these receptors will be also internalized and will satisfy this requirement. A number of studies where liposomes or lipid complexes were targeted to vitamin and growth factor receptors on cancer cells is given in Table I. Folate receptor is the only vitamin receptor with reported use in liposome targeting, although the use of other vitamin receptors such as that for pyridoxal phosphate or pyridoxine have been mentioned (Zalipsky et al., 1998). Growth factor receptors reported in liposome

TABLE I
LIGANDOLIPOSOMES TARGETED TO VITAMIN AND GROWTH FACTOR RECEPTORS

	Liposome composition	Substances delivered	Targeting ligand	Cell lines	Reference
FR	DSPC:Chol	Doxorubicin; marker	Folic acid	KB (human nasopharyngeal carcinoma), HeLa (human cervical	(Lee and Low, 1994, 1995); Wang and Low, (1998)
	HSPC:Chol:PEG-	Doxorubicin;	Folic acid	carcinoma) KB	Gabizon et al. (1999)
	DSPE Diplasmenyl(C_{16})	marker Ara-C; marker	Folic acid	КВ	Rui et al. (1998)
	choline: dihydroChol DSPC:Chol Anionic liposome	Oligonucleotides plasmid DNA– polylysine	Folic acid Folic acid	KB KB	Wang et al. (1995) Lee and Huang (1996)
HER-2/neu	PC:Chol; PC:Chol:PEG- DSPE (PC = POPC, HSPC, DSPC)	complex Doxorubicin; marker	Anti-HER2 Fab' Anti-HER2 scFv	SKBR-3, MCF-7, MCF/7HER2; MBA-MD-453; BT-474 (human breast carcinomas); WI-38 (normal	Kirpotin et al. (1997a, 1997b, 1998, 2000); Park et al. (1995, 199 1998a, 1998b, 2000)
	HSPC:Chol:PEG- DSPE	Doxorubicin, marker	Anti-HER2 IgG	human lung) N-87 (human gastric carcinoma)	Goren et al. (1996)

		Phosphorothiate	Anti-HER2 Fab'	SKBR-3	Meyer et al. (1998)
	DOTAP:DOPE:PEG- DSPE DDAB:Chol;	oligonicleotide Plasmid DNA	Anti-HER2 Fab'	SKBR-3	Park et al. (1997)
	DDAB:Chol:PEG- DSPE DPPC:Chol	[³ H]-inulin	Human EGF	Fibroblasts	Ishii et al. (1989)
EGFRr	DSPC:Chol:PEG- DSPE	(marker) ¹²⁵ I-labeled antibody	Anti-human EGFR IgG (C225)	DU-145 (human prostate carcinoma	Harding et al. (1997)
	DC-Chol:DOPE	Plasmid DNA	Human EGF	HEC-1A (adenocarcinoma) GCH-1 (human	Kikuchi <i>et al</i> . (1996)
VEGFR	DSPC:Chol	Anti-VEGF nucleic acid	Anti-VEGF nucleic acid	chorionic carcinoma) HUVEC (human vascular	Willis et al. (1998)
NGFR	EggPC:DPPE:Chol	aptamers FITC-dextran	aptamers Mouse NGF	endothelium) PC12 (pheochromo- cytoma), HS294 (human melanoma)	Rosenberg et al. (1987)

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such as folic acid, small peptides/proteins such as the native growth receptors, including antibodies or antibody fragments, small molecules antibodies against appropriate receptors example, heregulin, EGF, NGF, and VEGF) or vitamin receptors with to target growth factor receptors with their small-peptide ligands (for example, it is also possible and maybe even desirable in some instances low-molecular-weight ligands, these approaches are not exclusive. For antibody fragments, and the targeting of vitamin receptors with their factor receptors has been primarily accomplished with antibodies or ing. It should be noted, however, that although the targeting of growth the following sections in comparison to other modes of liposome target small effector molecules or an antibody and are discussed at length in ogy in liposome targeting to internalizable cell surface epitopes using approaches make excellent case studies outlining general methodol-HER2 Fab' or single-chain Fv-targeted liposomes (Table I). These two folic acid tethered to the end of a PEG-linked lipid anchor and the antiapproaches are the folate-targeted liposomes using the small-molecule factors, and nucleic acid aptamers. The two most well studied of these (Table I). A variety of different ligands have been used to target these lar endothelial growth factor receptor, and nerve growth factor recepton targeting studies include the epidermal growth factor receptor, vascu-

II. TARGETING OF LIPOSOMES TO VITAMIN RECEPTORS: THE CASE OF FOLIC ACID

A. FOLATE RECEPTOR AS INTERNALIZING TARGET ON MALIGNANT CELLS

The vitamin folic acid and its reduced derivatives can be taken up by cells using both a low-affinity ($K_d \sim 1-5 \,\mu$ M) transmembrane protein responsible for the passive diffusion of reduced folates across the plasma membrane of cells and a high-affinity GPI-anchored protein ($K_d \sim 0.01-1 \,\mathrm{nM}$) that accumulates folic acid in the cell following receptor-mediated endocytosis by a clathrin-independent pathway (Kane et al., 1986, 1989; Kamen et al., 1988). The former is referred to as the reduced folate carrier (RFC) and demonstrates a considerable preference for reduced folates, such as 5-methyltetrahydrofolate, methotrexate, and 5-formyltetrahydrofolate, compared to free folic acid (Henderson, 1990; Antony, 1992). The RFC is unable to bind or mediate the uptake of FA conjugates. The folate-binding protein (FBP), also referred to as the folate receptor (FR), is actually a class of receptors expressed in low levels on some normal epithelial cells (FR- α Zimmerman, 1990;

(nystatin) or internalization (phorbol-12-myristate) were shown to inhibit uptake of FA conjugates (Lee et al., 1996), although neither drug Boerman et al., 1991; Weitman et al., 1992, 1994; Garin-Chesa et al., overexpressed in a number of different cancers (Mattes et al., 1990; 1999), and placenta (FR- β Ratnam et $\acute{a}l$., 1989). However, it is highly hematological cells (FR-β and FR-γ; Shen et al., 1994; Reddy et al., Holm et al., 1991, 1992, 1993; Ross et al., 1994; Patrick et al., 1997), is overexpressed in hematological cancers (Ross et al., 1994). A wide vaoverexpressed in carcinomas such as ovarian carcinomas, while FR- $\!\beta$ various conjugates are taken up by cancer cells by receptor-mediated endocytosis (Antony $et\ al.$, 1985; Leamon and Low, 1991; Turek $et\ al.$, in folate-mediated targeting to malignant cells and tissues expressing riety of different therapeutic and diagnostic agents have been studied 1993; Ross et al., 1994; Toffoli et al., 1997). For the most part, FR-lpha is reviewed (Reddy and Low,1998; Wang and Low, 1998). Folate and its Mathias et al., 1996, 1998; Lu et al., 1999). These studies were recently FR (Leamon and Low, 1991; Leamon et al., 1993; Lee and Low, 1995; the FR becomes localized to non-clathrin-coated pits known as caveoto some extent still controversial. It is has been demonstrated that other studies, folate receptors appear not to colocalize with caveolae mode of internalization, two specific inhibitors of caveolae assembly lae (Rothberg et al., 1990; Turek et al., 1993). Lending support to this 1993; Lee and Low, 1994). The route of internalization for the FR is of caveolae in folate uptake, FR chimeras targeted to clathrin-coated inhibits clathrin-mediate endocytosis (Smart et al., 1994). However, in up by caveolae and resided in multivesicular bodies at early time points port 5-methyltetrahydrofolate into the cytoplasm (Ritter et al., 1995) pits rather than caveolae were found to be unable to efficiently trans-(Mayor et al., 1994; Wu et al., 1997). Lending further support to the role very high numbers of the receptor (Rijnboutt et al., 1996). A diagram docytosis may be a relatively minor pathway for cells that overexpress Turek et al. (1993) showed that folic acid conjugates of BSA were taken indicating the various possible routes of uptake for both vitamin and transferrin conjugates at later times. However, clathrin-mediated en-(<60 min), but converged with the clathrin-mediate pathway used by molecule and the receptor, where cross-linking of the receptors leads to growth factor receptors is given in Fig. 2. The final destination may be a definitive answer as to the complete route of internalization following targeting to caveolae. Despite the large volume of work on the subject the folic acid conjugated to the liposome surface, liposomes may favor uptake into caveolae (Mayor $\it et~al.,~1994$). Due to the multiple copies of dependent on the multiplicity of the interaction between the targeted

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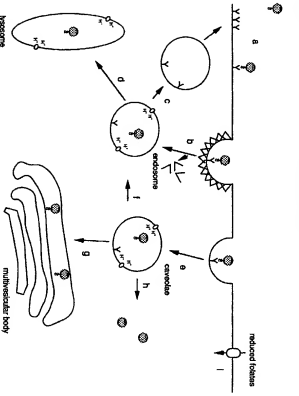


Fig. 2. Potential fates of ligand-targeted liposomes following binding to a target cell. Upon binding to cell-surface receptors, liposomes can either remain bound at the cell-surface, disassociate from the receptor, or accumulate in coated or noncoated invaginations. Following clathrin-mediated endocytosis (a), liposomes can be delivered to lysosomes (c) where they and their contents may be degraded by lysosomal peptidases and hydrolases. Receptors may be recycled back to the cell surface (b) or targeted for degradation in the lysosome (c). Some GPI-anchored receptors, such as the FR, can be taken up by caveolae-mediated endocytosis. Following internalization, the conjugated molecule can remain in caveolae, be transported to multivesicular bodies (g) or into the cytoplasm (h) and possibly reenter the lysosomal directed pathway (f) A substantial proportion of the folate-targeted molecules or liposomes appear to remain in a nondegradative compartment, allowing greater feasibility for delivering labile molecules via this route.

receptor binding is not known. An answer to this question may prove very important when selecting various molecules for encapsulation or complexation to lipid-based carriers.

Since FR- α is a GPI-anchored protein, phospholipase C treatment of FR- α -overexpressing cells results in a considerable loss of uptake of FA conjugates (Leamon and Low, 1992). In addition to the folate receptor, other receptors commonly found in caveolae include receptors for platelet-derived growth factor (PDGF receptor) (Liu et al., 1996), bradykinin (de Weerd and Leeb-Lundberg, 1997), insulin (Goldberg et al., 1987), and epidermal growth factor (EGFR; Mineo and Anderson, 1996). An extremely important advantage of targeting to this endocytic

pathway as compared to clathrin-mediated endocytic pathway is that the final destination of the targeted molecules in clathrin-mediated endocytosis is the lysosome, where degradative enzymes can degrade labile drugs, genes, or other biomolecules. The caveolae pathway appears to be rather nondestructive, as molecules are able to remain intact for up to days following binding to internalizing receptors (Leamon and Low, 1991; Wang et al., 1995). This results in a significant enhancement of the activity for the delivered molecule (Leamon et al., 1992; Rui et al., 1998). In addition, similar to clathrin-coated vesicles, caveolae are also acidified to a relatively low pH (Lee et al., 1996), allowing for the development of pH-sensitive liposomes for enhanced cytoplasmic delivery (Lee and Huang, 1996; Reddy and Low, 1998; Rui et al., 1998).

B. FOLIC ACID AS A TARGETING LIGAND: COUPLING TO LIPOSOMES

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posome in a way that does not impair receptor-binding properties of molecules must be stably conjugated on the outer surface of the li-Low, 1994). In order to act as liposome-targeting ligands folic acid ing their natural ligand, folic acid, as a targeting moiety (Lee and with FR. Ligands are attached to liposomes by conjugation to lipid FA and allows unhindered interaction of the liposome-conjugated FA erol derivatives, satisfy this requirement. Opposite to this hydrophotwo closely positioned C₁₆-C₂₀ alkyl or acyl chains, such as diacylglyc molecules sufficiently hydrophobic to act as "membrane anchors" staythan in the aqueous environment outside the liposome. Anchors with ing in the environment of hydrocarbon chains of the lipid bilayer rather gations. These lipid anchors are included in the liposome lipid compo N-derivatives are molecules of choice for many liposome-ligand conjutive group, such as primary amino, carboxy-, or thiol group, that can bic domain, lipid anchor molecules usually have a chemically reacbranotropic conjugates that are later used to form ligandoliposomes sition before ligand conjugation or conjugated to ligands to form mem Allen et al., 1997). Distearoyl phosphatidylethanolamine and its form a stable bond with the ligand (for review, see Park et al., 1997c; preformed liposomes via a simple lipid anchor or to a lipid anchor with a reactive conjugates, including liposomes. Folic acid has been coupled to out the loss of FR binding and has been used for making various FRdomain, of which the group in the γ-position can be modified with-The molecule of folic acid has two carboxyl groups in its glutamate PEG spacer having terminal primary amino group using water-soluble Targeting of liposomes to folate receptors was demonstrated by us-

aqueous solutions, and also eliminated the exposure of drug-loaded liand extrusion through nanoporous track-etched membranes to produce solvent; following the evaporation of the solvent, they were hydrated 2.8 vs 4.5, respectively) (Fan et al., 1991); however, the presence of can be separated by ion exchange chromatography due to the differ in the α -position and is not reactive with FR. These two components γ -carboxyl group; the remaining conjugate, however, contains the linker strategies is shown in Fig. 3. In this method, folate is first conjugated posomes to potentially deleterious coupling reagents and conditions coupling methods using chemically activated hydrophobic linkers ir process, avoided the difficulty of developing rigorous and reproducible effort. The liposome lipids and the conjugate were mixed in an organic protected glutamic acid, and then, pteroic acid to the activated ceptable, the conjugate can be constructed by stepwise addition of first monoderivatized PEG was isolated and reacted with the hydrophobi ing properties of subsequently produced FA-conjugated liposomes. The the α-derivative does not seem to have negative effect on the target formation of the liposomes (Lee and Low, 1995). One of these coupling carbodiimide EDAC (Lee and Low, 1994). In another approach, the con even permeabilization (Uster et al., 1996). Indeed, "anchor" N-succinyl-distearoylphosphatidylethanolamine using a sim In this reaction 70-80% of the conjugate contains the linker at the jugate was first synthesized and admixed into the lipids prior to the mediated targeting, compared to a protein such as an antibody or anti bic domains into the liposome bilayers without liposome destruction or modified poly(ethylene glycols) to merge spontaneously their hydropho tion of folic acid and other water-soluble vitamins to preformed, drug Finally, one may envision even more elegant methods for the conjuga together in an aqueous buffer and subjected to usual steps of dispersion PEG linker (Knepper et al., 1990), but at a higher cost and with more body fragment, is the relatively higher shelf life of the conjugate. Pro using a small, naturally occurring molecule such as folic acid for ligano 1999) and antibody fragments (Kirpotin $\it et\,al., 2000$ b). One advantage o as well as tor even larger ligands such as antibodies (Ishida and Allen peptide- and oligosaccharide-linked liposomes (Zalipsky et al., loaded liposomes based on the remarkable ability of hydrophobically 100-nm unilamellar vesicles . This allowed for a more efficient coupling lar conjugation step. If the presence of α-conjugated folate is unac teins are often more sensitive to changes in environmental conditions "insertion" method was recently demonstrated for preparation of pKs between the α - and γ -carboxyl groups of the foliate (pk the usefulness of

Fig. 3. Synthetic scheme for coupling of folic acid to a lipid anchor (1,2-distearoyl-3-sn-phosphatidylcholine; DSPE) via a poly(ethylene glycol) linker. Folic acid is first coupled to poly(ethylene glycol) bis-amine using dicyclohexylcarbodiimide as a coupling reagent. The monoderivatized folic acid-PEG conjugate is purified and subsequently coupled to N-succinyl-DSPE using a similar coupling reaction. N-succinyl-DSPE can be readily formed from DSPE and succinic anhydride in the presence of a weak organic base (Kung and Redemann, 1986).

and there may be significant stability concerns if they are stored for long periods at 4°C, conditions under which drug-loaded liposomes are normally stored.

C. Interaction of Folate-Targeted Liposomes with FR-Overexpressing Cells

of the cancer cells (Lee and Low, 1994). Little or no cell association overexpressing cells, they were quite capable of binding the milk folatewhile the liposomes with short-spacer FA conjugates did not bind to FR was seen when any of several short spacers were used to conjugate determined to be essential in distancing the targeting ligand from the acid to a lipid anchor already present in liposome membranes (Lee and folic acid to the surface of the liposome. However, we have noted that of the liposome to reduce steric hindrance to its receptor and to benepears important for the folic acid to be extended above the polymer coat chain of the more abundant unmodified conjugate (Fig. 4). Thus, it apand co-workers (1999) showed that the length of the spacer was also binding of the liposome-tethered folic acid to the cell surface receptor; bility of the liposome-ligand spacer was necessary to allow unhindered binding protein, which is essentially an extracellular domain of the foliposome surface to allow binding to the folate receptor on the surface allow its better access to cell membrane receptors. fit from the conformational flexibility of the polymer linker which may jugated to liposomes using a PEG spacer greater in length than the PEG lifetimes. Cell association was markedly increased when folate was containing also unmodified PEG-DSPE used to increase their circulation important when using FA-PEG-DSPE conjugates on the liposomes con the same may be true for other small-molecule ligands as well. Gabizon 1993, unpublished observation). Apparently, certain length and flexilate receptor shed from the cell membranes (Kirpotin and Kolhouse, Low, 1994, 1995). The PEG spacer, linking FA to the lipid anchor, was Folate-targeted liposomes were first prepared by conjugation of folio

Lee and Low (1994) studied the kinetics of folate–liposome associations with the cells. When targeted to FA- α -overexpressing KB cells, FA-derivatized liposomes were seen only at the cell periphery at early times, but were seen throughout the cytoplasm as a punctate fluorescence pattern at later times (4 h). Both the kinetics of internalization and the total number of liposomes bound at saturation (2.5 \times 10⁵) were lower than for the much smaller albumin conjugates studied in an earlier work (Leamon and Low, 1991). The authors suggested the latter may be a result of the multivalent nature of FA display when present

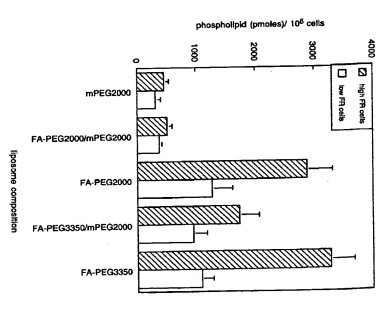


Fig. 4. Effect of PEG length on binding of FA-derivatized liposomes to KB cells. Liposome binding was determined by measuring the amount of cell associated [³H]Chol. Liposomes were incubated with KB cells having low (low FR) or high either low- or high (high FR)-folate receptor expression. High expression of folate was achieved by passaging the cells in folate-free medium (the only source of folate being endogenous folate in fetal calf serum, added to the medium at 10%). Liposomes contained folate-PEG-DSPE conjugates with PEG having molecular weights of either 2000 or 3350 (PEG2000 and PEG3350) and in either the presence or absence of PEG2000-DSPE at approximately 7% of total liposomal phospholipids. The cells were incubated with liposomes for 24 h at 37°C. From Gabizon et al. (1999), with permission from the American Chemical Society.

in the form of liposomes. The depressed rate of internalization may result from an increased steric hindrance to cell surface receptors due to the larger size of the liposome, compared to albumin. FA liposomes were also shown to be endocytosed by looking at the difference in acid-removable liposomes following incubations at both 4° and 37°C (Lee and Low, 1994). While FA liposomes incubated with KB cells at 4°C could be quantitatively removed from the cells by an acid wash, only

LIPOSOME TARGETING TO TUMORS

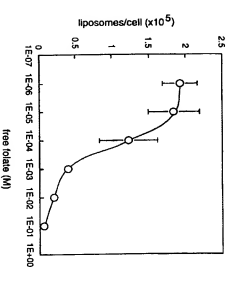


Fig. 5. Effect of folic acid on the uptake of folate-conjugated liposomes by KB cells. KB cells were incubated with calcein-loaded FA-derivatized liposomes for 4 h at 37° C in the presence of varying concentrations of free FA. The amount of cell-associated liposomes was determined from the calcein fluorescence in the detergent lysates of thoroughly rinsed cells. From Lee and Low(1994) with permission from ASBMB.

half of the liposomes could be removed following incubation at 37°C for 4 h. It is interesting that there are still a relatively high number of liposomes at the cell surface even at 4 h, a time where other receptors have been shown to be almost completely internalized to an intracellular localization. This may be a result of the different trafficking pathways involved in caveolae-mediated endocytosis compared to the more typically studied clathrin-mediated endocytosis, which terminates in the lysosome.

This association was shown to be specific for folate-mediated targeting by competitive inhibition studies with either free FA or antiserum against FR- α (Lee and Low, 1994). The inhibition of FA-derivatized liposome association with KB cells by free FA is shown in Fig. 5. Interestingly, a concentration approaching 1 mM free FA was required to inhibit FA-derivatized liposome uptake by KB cells (Fig. 5). This is in contrast to the relatively low $K_{\rm d}$ of free FA for the FR (0.01–1 nM; Kamen et al., 1988) and the significantly lower concentrations of free FA (<200 nM) required to inhibit uptake of FA-deferoxamine conjugates (Wang et al., 1996). The multivalent binding of liposome bound conjugates likely leads to this reduced inhibition and gives rise to a targetable therapeutic that may be useful in vivo when considering the

relatively low free FA concentrations in plasma (<20 nM; Antony et al. 1985; Kamen et al., 1988; Antony, 1992).

D. Delivery of Antineoplastic Drugs to Cancer Cells

D. Delivery of Antineoplastic Drugs to Cancer Cells by Folate-Targeted Liposomes

centuate the favorable results seen in cell-culture studies even further cytotoxicity in cell culture studies, the area under the concentration veret al., 1999). For example, although there is only a 1.6-fold increase in promising results due in part to the favorable pharmacokinetic propertrapolation of these results to in vivo conditions gives rise to even more overexpressing KB cells when compared to both free doxorubicin (1.6liposomes was shown to increase the cytotoxicity of doxorubicin to FR 1995; Rui et al., 1998). Folate-mediated targeting of doxorubicin-loaded targeted liposomes to FR-overexpressing cancer cells (Lee and Low sus time curve for tumors is approximately 8-fold greater for liposomal ties of liposomal carriers (Hwang, 1987; Allen et al., 1995a; Drummond fold) and nontargeted liposomal doxorubicin (45-fold; Fig. 6). The exexposure of the cancer cells to the drug in vivo when encapsulated in doxorubicin compared to free doxorubicin, meaning there is a greater liposomes (Unezaki et al., 1995; Gabizon et al., 1997). This should ac-Several studies have reported the cytotoxicity of drug-loaded FA

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Ara-C has been effectively delivered to KB cells using FA targeting of pH-degradable liposomes composed of di-C₁₆-plasmenylcholine (Rui et al., 1998). The IC₅₀ of ara-C was increased approximately 6000-fold when delivered in folate-targeted pH-degradable liposomes (0.49 µM) compared to free ara-C (2.8 mM). These liposomes are programmed to decompose at a pH characteristic of endosomal compartments. While the authors suggest the heightened sensitivity to ara-C is due in large part to the pH-sensitive lipid composition, it is also possible that targeting to what has already been shown to be a relatively nondestructive endocytic pathway contributes substantially to the increased cytotoxicity. This approach may prove especially fruitful for delivering readily degradable drugs or macromolecules.

E. Delivery of Genes and Antisense Oligonucleotides to Cells Using Folate Targeting

Folic acid has also been used as a ligand to target genetic material and antisense oligonucleotides to FR-overexpressing cells (Gottschalk *et al.*, 1994; Wang *et al.*, 1995; Lee and Huang, 1996; Reddy and Low, 1998) Folic acid-derivatized polylysine conjugates were used to condense DNA

and antisense oligonucleotides and deliver them specifically to tumor cells (Gottschalk et al., 1994; Citro et al., 1992, 1994; Ginobbi et al.,

be dependent on the conjugated folate and was also markedly enhanced

1997). Gene expression using these targeted complexes was shown to

when coincubated with a replication-defective adenovirus, indicating

the need for endosomal disruption for efficient release of DNA from

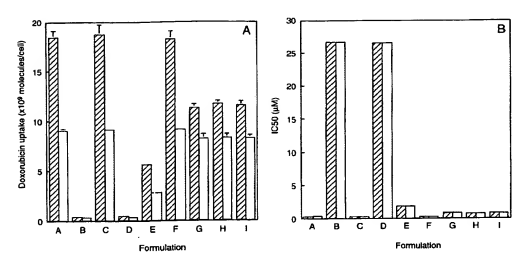


Fig. 6. Cellular uptake and cytotoxicity of folate-targeted and nontargeted formulations of doxorubicin. KB cells (cross-hatched bars) or HeLa cells (open bars) were incubated with doxorubicin formulation (100 µM DOX): folate-conjugated liposomal DOX without PEG coating (FA-L-DOX) (A); DOX in nontargeted, noncoated liposomes (L-DOX) (B); folate-conjugated PEG-coated liposomal DOX (FA-SSL-DOX) (C); nontargeted PEG-coated liposomal DOX (SSL-DOX) (D); FA-L-DOX + 1 mM folic acid (E); FA-L-DOX + 20 nM 5-methyltetrahydrofolate (F); free DOX (G); free DOX + "empty" folate-conjugated liposomes (H); free DOX + "empty" nontargeted liposomes (I). Cell-associated doxorubicin was determined by fluorescence spectroscopy, and cytotoxicity was determined using a tetrazolium (MTT) assay. IC₅₀ refers to the concentration of doxorubicin that results in 50% decrease in viability compared to sham-treated control. Adapted from Lee and Low (1995), with permission from Elsevier Science.

expression was independent of folate targeting. However, at higher raposomes (Drummond and Daleke, 1997). anionic liposomes containing the "caged" pH-sensitive lipid N-citracomplexes (DC-Chol:DOPE) and were significantly less cytotoxic. A simtively small size (\sim 74 nm), folate targeting resulted in a 20- to 30-fold tios, where the complexes obtained a net negative charge and a relacondensed DNA ratios the complexes were positively charged and gene cle composed of polylysine-condensed DNA, DOPE, and the pH-sensitive intracellular organelles (Gottschalk *et al.*, 1994). conyldioleoylphosphatidylethanolamine (N-cit-DOPE; Reddy and Low increase in transfection efficiency when compared to cationic lipid–DNA lipid cholesteryl hemisuccinate (CHEMS). At low lipid-to-polylysine cles may enhance transfection efficiencies even further, similar to the ilar approach complexed polylysine-condensed DNA with FA-targeted Huang (1996). Other folate-targeted cationic lipid based delivery vehi-4 to 5 when compared to the similar composition described by Lee and Daleke (1995) and later shown to be useful in preparing pH-sensitive li 1998). This "caged" lipid was originally described by Drummond and $N ext{-cit-DOPE}$ was shown to increase transfection efficiency by a factor of Lee and Huang (1996) developed a pH-sensitive lipid-DNA parti-The use of this "caged'

Antisense oligonucleotides against the EGFR were delivered to KB cells via folate targeting to cultured KB cells (Wang et al., 1995). The liposomes utilized in these experiments were composed of eggPC, cholesterol, and either with or without folate-PEG-DSPE. A fluoresceinlabeled oligonucleotide was used to show a 16-fold increase in uptake over free oligonucleotides and a 9-fold increase over nontargeted liposomal oligonucleotide. Uptake could be inhibited by free folic acid (1 mM), demonstrating the specificity of internalization. However, delivery to the nucleus was relatively inefficient, likely due to the relatively stable lipid composition used in this particular formulation and thus inability to escape the confines of the endosomes or lysosomes. Finally, while folate-specific growth inhibition did occur, there was no significant difference in growth inhibition between the anti-EGFR oligonucleotide

via receptors, such as the folate receptor.

results seen with anti-HER2 targeted cationic lipid-DNA complexes

(Section III,E). For lipid-based gene delivery vehicles, active targeting

LIPOSOME TARGETING TO TUMORS

and the scrambled sequence. These results suggest that folate targeting may be effective at delivering both oligonucleotides and plasmid DNA to receptor-overexpressing cells. However, the efficiency of delivery, both from the endosome and under *in vivo* conditions, needs to be increased and studied further.

F. IN VIVO IMPLICATIONS FOR FOLATE-MEDIATED LIPOSOME TARGETING

To date, folate targeting of liposomes has been almost exclusively studied in cell culture, with minimal work being completed in vivo. Mathias and co-workers (1996, 1998, 1999) have used folic acid to target small-molecule radiotracers for imaging folate receptor-overexpressing tumors in vivo. However, the pharmacokinetics of small-molecular-weight compounds such as these are undoubtedly different from those of sterically stabilized or even conventional liposomal carriers. For example, these relatively small-molecular-weight radioconjugates are found in high amounts in the kidney, an organ that would be relatively inaccessible to large liposomal carriers approximating 100 nm in size.

An important concern for use of a folate receptor targeting approach in vivo involves the toxicity to normal healthy tissues, specifically those of hematopoietic lineage. The β -isoform of the folate receptor (FR- β) is expressed in high levels on hematopoietic cells (Ross *et al.*, 1994; Shen *et al.*, 1994; Reddy *et al.*, 1999), and thus targeting of cytotoxic agents to this receptor may be expected to result in significant bone marrow toxicity. Fortunately, Reddy and co-workers (1999) were able to demonstrate that CD34+ hematopoietic cells do not bind FA-conjugated liposomes, although these cells overexpress FR- β .

The final potential problem with targeting to cell-surface receptors using FA-conjugated liposomes is the binding-site barrier phenomenon observed for tumor-specific antibodies (Juweid $et\ al.$, 1992; Fujimori $et\ al.$, 1990): the binding of FA conjugates to FR- α at the site of liposome extravasation may limit its penetration of the tumor and thus accessibility to cancerous cells located any significant distance from the supporting vasculature. However, limited diffusion and the increased bioavailability of encapsulated drugs may be enough in itself to provide a significant improvement to nontargeted liposomes. An alternative approach may be to use antibody fragments against the folate receptor rather than use the ligand itself. The reduced avidity of the targeting ligand for the receptor may result in a more even tumor distribution of the carrier, similar to that seen with anti-HER2-targeted immunoliposomes (Section III,D). While speculation on the feasibility of targeting FRs in vivo suggests both significant promise and a degree of

uncertainty, the true test lies in the completion of efficacy studies in appropriate animal tumor models, such as in the treatment of ovarian cancers.

G. OTHER VITAMIN RECEPTORS AS TARGETS FOR LIGANDOLIPOSOMES

tion (Wilbur et al., 1999; McEwan et al., 1999). gies will likely be attempted. At present, pyridoxine, pyridoxal phosceptors of these vitamins become better understood with respect to strategies can be easily adapted for conjugation of these ligands to lipomalignant cells, further development of other vitamin-targeted strateroutes of internalization and expression patterns on both normal and the use of other vitamins as ligands for liposomal delivery. As the revelopment of folate targeting laid the methodological groundwork for receptors for vitamins other than folate. However, the successful dereported, and the chemistries appears amenable to liposome conjugasomes. Preparation of fully functional conjugates of cyanocobalamine geting ligands other than liposomes for therapeutic or diagnostic agents phate, biotin, riboflavin, and nicotinamide have been reported as tar-(B₁₂) with radioiodinated markers and spacer molecules was recently (Low et al., 1997a,b; Holladay et al., 1999). The developed conjugation Few data are available on the targeting of liposomes to cells using

Aside from specific targeting of their receptors, vitamins have been used in distinct roles in the development of liposome technology. Thus, biotin has been used as an adaptor molecule for conjugating targeting ligands to liposomes via a noncovalent biotin-streptavidin linkage (Rosenberg et al., 1987; Harasym et al., 1995; Wong et al., 1997). o. Tocopherol (vitamin E) has been used to prevent oxidative damage to liposomes during storage (Barenholz et al., 1993). These functions of vitamins in liposome technology are out of the scope of this chapter. However, it is safe to say that the groundwork for the use of vitamins as liposome-targeting ligands in the clinic has been firmly set and awaits a promising future.

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III. TARGETING LIPOSOMES TO GROWTH FACTOR RECEPTORS

A. GROWTH FACTOR RECEPTORS AS RECOGNITION MOLECULES OF MALIGNANT CELLS

Like their effectors, hormone receptors present a family of structurally and functionally different proteins. Some hormone receptors, such as steroid receptors, are intracellular proteins that are inaccessible

endocytosed by receptor-bearing cells following binding of their approreceptors are attractive targets due not only to their high expression Chrysogelos and Dickson, 1994; Fix, 1994; Press et al., 1997). These poor survival (Berchuck et al., 1990; Borg et al., 1990; Toi et al., 1991; of these receptors are associated with an increased risk of recurrence or (Yu et al., 1991, 1994; Niehans et al., 1993). Often, high expression levels receptors have been associated with tumor proliferation or metastasis et al., 1991; De Vries et al., 1992). The expression levels of these various dothelial growth factor (VEGF) receptors (Shibuya et al., 1990; Tischer cular endothelial cells supporting the tumor overexpress vascular enet al., 1987, 1989; Mansson et al., 1989; Beitz et al., 1992; Khazaie et al., expressed in different malignancies (Sporn and Roberts, 1985; Slamon (FGF receptor), and HER2/neu receptor have all been shown to be overmal growth factor receptor (EGFR), fibroblast growth factor receptor that prove most valuable for cell-targeted delivery of drugs. The epidermains are exposed on the cell surface. It is these accessible receptors membrane, are plasma membrane proteins whose effector-binding doas peptide hormones and growth factors that do not permeate the cel what can be called a "Trojan horse" approach. tially allows the therapeutic agent access to an internal localization in receptors discussed above, targeting to one of these receptors poten-Hurwitz et al., 1995; Kirpotin et al., 1997a). Thus, similarly to vitamin priate ligand or an antibody agonist (Pastan and Willingham, 1981; levels on malignant cells but also because they have been shown to be 1993; Fox et al., 1994; Scher et al., 1995). In addition, angiogenic vasfusion across biological membranes. Receptors for other hormones, such for targeting. Steroid hormones may reach their receptors following dif Tagliabue et al., 1991; Sorkin and Waters, 1993; French et al., 1994;

The HER2/neu receptor recently became an extensively studied target for liposomes and lipid-based gene- and drug-delivery systems. This receptor is a glycosylated transmembrane protein of approximately 185 kDa that possesses tyrosine kinase activity. It is a member of the epidermal growth factor receptor (EGFR) family and is involved in many growth-signaling pathways within the cell, alone or in cooperation with other members of EGFR family. Several excellent reviews have described the basic biology and biochemistry of the HER2/neu receptor (Hynes and Stern, 1994; Hung and Lau, 1999). This protein has high levels of overexpression in different malignancies (10⁵–10⁶ receptors/cell), low level of expression in healthy tissues, and homogeneous and stable expression in primary tumors and sites of metastasis (Press et al., 1990; Lewis et al., 1993; Niehans et al., 1993). These characteristics made this protein a prime molecular target for cancer

immunotherapy using a recombinant humanized monoclonal antibody (trastuzumab) and has recently made its way into the clinic (Baselga et al., 1999; Pegram et al., 1998; Shak, 1999). These properties are also likely responsible for the many encouraging preclinical results obtained with an anti-HER2-targeted immunoliposome carrying encapsulated doxorubicin (Section III,D).

B. Design of HER2-Targeted Immunoliposomes

In contrast to the folate receptor targeting described above, where the natural ligand for FR was also the liposome-targeting ligand, targeting of liposomes to HER2 receptor exemplifies a different approach in which the targeting ligand is an antibody against extracellular portion of the receptor.

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receptors and especially HER2/neu satisfy the criteria for target antivehicles (Table II). As discussed in the preceding section, growth factor successful practical design of immunoliposomes as cancer drug-delivery made their way to the clinic, one can put forward a set of criteria for a munoliposomes or, in fact, any ligand-targeted liposomes have not yet compasses two decades of research (Allen et al., 1997). Although imliposome requires only the presence of the antigen-binding domain, leaving a researcher with a choice of liposome-conjugated antibodies. gen selection quite well. Evidently, the targeting function of an immunoenhanced blood clearance of immunoglobulin-conjugated liposomes me A popular option of conjugating whole immunoglobulin molecules to association of immunoliposomes with target cells while preserving long-circulating properties (Shahinian and Silvius, 1995; Zalipsky et al., tion of immunoliposomes is desirable in order to prolong their circu maleimide chemistry (Martin et al., 1981; Martin and Papahadjopoulos, at the hinge region provided a unique conjugation site by efficient thiol HER2 Fab' fragments proved to be a better solution. Cysteine residues diated by mononuclear phagocyte Fc receptor (Aragnol and Leserman. tion site, immunogenicity of xenogeneic IgG (Harding et al., 1997), and liposomes appears suboptimal because of the poorly defined conjugaof Fab' fragments to the maleimide-activated terminal end of a lipoders the binding of an antibody to cell surface receptors if the spaces 1986; Derksen et al., 1988). The use of recombinant humanized antisome-linked PEG-lipid (PEG-DSPE) (Fig. 7) resulted in an unimpeded between the antibody and the lipid anchor is too short. Attachment lation lifetime for better distribution into a tumor, PEG coating hin 1982). Similarly to folate-conjugated liposomes, when steric stabiliza The history of antibody-targeted liposomes (immunoliposomes) en-

sign Criteria for Ligand-Directed Liposome Targeting

DES	DESIGN CRITERIA FOR LAGANU-DIRECTED LAFOSOME LARGETING
Component	Considerations for optimum design
Target antigen	Expression Highly and homogeneously overexpressed in target tissue
	Function Vital to tumor progression, so that down-modulation does not occur or is associated with therapeutic benefit
	Shedding of antigen Limited, to avoid binding to soluble antigen and accelerated clearance
Targeting ligand	Affinity

Tinity
High enough to ensure binding at low liposome concentrations
Low enough to avoid "binding-site barrier" effect (Weinstein)

Immunogenicity

Humanized MAb, to remove murine sequences. Use fragments without Fc portion (Fab', scFv) to avoid interaction with Fc recentor

Small molecular weight ligands should not be immunogenic—may act as haptens

Internalization

Efficient endocytosis by target cells is desirable

Production

Fear and econo

Easy and economical scale-up, e.g., by efficient bacterial expression system

Stable during storage

Ligand-liposome linkage

Stability

Covalent attachment to hydrophobic anchor, stable in blood

Attachment site

Away from the binding site to ensure correct orientation of antibody or ligand molecule.

Well-defined, to ensure reproducibility and uniformity of coupling
Avoids steric hindrance (e.g., from PEG) of ligand binding a

Avoids steric hindrance (e.g., from PEG) of ligand binding and internalization

Chemical nature of the linker

Nontoxic, nonimmunogenic, and avoids opsonization
Does not affect drug loading or membrane stability
Excess linker may be quenched to avoid nonspecific coupling to
biomolecules

Good availability, economical manufacturing process

Liposome

١.,

Stable as intact construct in vivo

(continued)

Table II (continued)

Component	
Considerations for optimum design	

Pharmacokinetics
Long circulating
Tumor penetration
Capable of extravasation in tumors
Small diameter improves penetration
Encapsulation

Small diameter improves penetration into tumor tissue Encapsulation Efficient, high capacity (e.g., by remote loading)

Drug

Encapsulated drug storage-stable and resists leakage

Bystander effect

Drug affects tumor cells not directly targeted (bystander cells)

Interaction with target cells
Effective against target cell population
Cytotoxicity enhanced by binding of ligand

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sequences and thus are likely to have little or no immunogenicity in from phage-display libraries (Marks et al., 1992; Schier and Marks, targeting antibody may be single-chain Fv (scFv) fragments selected a C-terminal cysteine group, away from the antigen-binding site. Last component of liposome targeting (Nielsen et al., 2000). phage internalization into live target cells (Becerril et al., 1999) allows but not least, a recently reported scFv selection technique based on engineered to contain a unique liposome conjugation site, for example, produced in quantities in bacterial hosts, are easily purified, and can be Fvs are relatively small (26–29 kDa) recombinant proteins that can be broad range of specificities and binding characteristics. Single-chain Marks, 1996; Schier et al., 1996) allow construction of scFvs with a humans. Phage-display selection and affinity maturation (Schier and 1996). Single-chain Fvs are constructed from human immunoglobulin 1996, 1998; Kir $potin\ et\ al.,\ 1998).$ An even better choice of liposomefor selective internalization by the target cells, which is a desirable the creation of scFvs optimized not only for selective binding, but also

Anti-HER2 immunoliposomes targeted to HER2-overexpressing tumors were a subject of several recent studies (Goren et al., 1996; Park et al., 1995, 1997a, 1997b, 2000; Kirpotin et al., 1997a, 1997b, 1998, 2000a). In these studies anti-HER2 immunoliposomes were constructed on the platform of sterically stabilized PEG-coated liposomes containing doxorubicin encapsulated by the ammonium ion gradient method. These liposomes are similar to liposomal doxorubicin, which has been recently introduced into the clinic under the trade name of Doxil (Alza Corp.) (Martin, 1998; Gabizon and Barenholz, 1999).

Fig. 7. Synthetic scheme for conjugation of Fab' fragments to liposomes. An amino-PEG-DSPE conjugate is derivatized with the bifunctional cross-linking reagent, N-(γ-maleimidopropionyloxy)-succinimide The obtained maleimide-terminated PEG-DSPE derivative can be incorporated into liposomes and subsequently derivatized with a Fab' or scFv fragment via reduced sulfhydryl group of peptide terminal cysteine, located away from the antigen-binding site specific for a cell-surface receptor. Alternatively, the fragments can be conjugated with maleimido-PEG-DSPE in solution and then captured into the liposome bilayer by co-incubation of the liposomes with the conjugate (Kirpotin et al., 2000a).

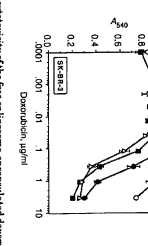
Fig. 8. Uptake of rhodamine-labeled antiHER2 immunoliposomes (left column) and fluorescein-labeled transferrin (middle column) by HER2-overexpressing SKBR3 cells (A,B) or by MCF-7 cells having low expression of HER2 (C). The cells were incubated with liposomes at 0.1 mM of liposome phospholipid and 37°C for 10 min (A) or 30 min (B and C). Liposome localization was visualized by confocal microscopy of intact cells. Superimposed images are shown in the right column. From Kirpotin et al. (1997) with permission from the American Chemical Society.

C. IN VITRO STUDIES WITH ANTI-HER2 IMMUNOLIPOSOMES

Interaction of anti-HER2 sterically stabilized immunoliposomes with target cancer cells was first studied in the cultures of human breast carcinoma cells of either low (MCF-7; 10⁴ receptors/cell) or high (SKBR-3) 10⁶ receptors/cell) expression levels of the HER2 receptor. Confocal fluorescence microscopy showed colocalization of anti-HER2 targeted im-

Interaction of anti-HER2 sterically stabilized immunoliposomes with target cancer cells was first studied in the cultures of human breast carcinoma cells of either low (MCF-7; 10⁴ receptors/cell) or high (SKBR-3; 10⁶ receptors/cell) expression levels of the HER2 receptor. Confocal fluorescence microscopy showed colocalization of anti-HER2 targeted immunoliposomes and fluorescein-labeled transferrin (Fig. 8), indicating uptake by HER2-overexpressing cells of liposomes into the clathrin-mediated endocytic pathway (Park et al., 1995; Kirpotin et al., 1997a). MCF-7 cells expressing low levels of the HER2 receptor were able to

LIPOSOME TARGETING TO TUMORS



PEG coating ("conventional" immunoliposomes) (●), PEG-coated anti-HER2 immunocells. The cells were treated with doxorubicin in anti-HER2 immunoliposomes without free doxorubicin (□). From Park *et al.* (1995) with permission. © 1995 by the National liposomes (○), PEG-coated immunoliposomes conjugated to an irrelevant Fab' (■), or Fig. 10. In vitro cytotoxicity of the free or liposome-encapsulated doxorubicin in SKBR3

tial experiments, the antibody was conjugated directly to the liposome anti-HER2 immunoliposomes (Park et al., 1995; Fig. 10). In these iniconjugated to a HER2-specific antibody that did not induce internalizapresumably due to an interference of receptor binding by the conjugated were shown to also reduce the cytotoxicity of the targeted formulation doxorubicin. Liposomes containing the sterically hindering PEG-DSPE significantly less toxic than either anti-HER2 immunoliposomes or free somes or liposomes containing an irrelevant antibody were shown to be SKBR-3 cells as free doxorubicin (IC₅₀ = $0.3 \mu g/ml$). Nontargeted lipomunoliposomes were shown to be as cytotoxic to HER2-overexpressing surface rather than to the distal end of a PEG spacer. These targeted impolymer. Targeted immunoliposomes were also relatively noncytotoxic creasing the effectiveness of doxorubicin delivery into target cells (Goren et al., 1996). This indicates how essential endocytosis is to intion were only as cytotoxic as the nontargeted liposomal formulation liposomes to the HER2 receptor does not ensure endocytosis. Liposomes to HER2 negative lung fibroblast cells (WI-38). Binding of the immuno-Cytotoxicity experiments were performed with doxorubicin-loaded

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D. IN VIVO STUDIES WITH ANTI-HER2 IMMUNOLIPOSOMES

models have yielded very promising results using anti-HER2 immuno-In vivo pharmacokinetic and antitumor efficacy studies in xenograft

able to effectively deliver the oligonucleotide payload to the cell nuet al., 1998). In this case, not only were the targeted complexes en cationic lipids and complexed with antisense oligonucleotides (Meyer of the oligonucleotide. (Right column) Nontargeted complexes. From Meyer et al. (1998) PEG chains was conjugated to the anti-HER2 Fab' (left column) show nuclear delivery munoliposomes depended on the number of conjugated anti-HER2 Fab', clei (Fig. 9). Both binding and endocytosis of anti-HER2-targeted imdocytosed, but only those containing anti-HER2-conjugated Fab' were results were obtained with anti-HER2-targeted liposomes containing take up the transferrin conjugate but not the immunoliposomes. Similar by low-HER2-expressing MCF-7 cells was undetectable. conditions, more than 80% of the liposomes were endocytosed. Uptake 8,000-25,000 liposomes per cell after 3-4 h of incubation. Under these in the cell growth medium, liposome uptake reached a maximum of posome (Kirpotin et al., 1997a). At 0.025 mM of liposome phospholipid which reached a plateau at approximately 30-40 Fab' per 100 nm liwith permission from ASBMB.

oligonucleotide (PS ODN) in HER2-overexpressing SKBR3 cells. (Top row) FITC-labeled

liposome (DOTAP:DOPE:PEG-DSPE) complex with FITC-labeled phosphorothicate

Fig. 9. Intracellular distribution of the components of PEG-DSPE-stabilized cationic

PS ODN; (bottom row) rhodamine-labeled lipid. Only the complexes where a portion of

liposomes targeted by anti-HER2 Fab' and scFv (Park et al., 1997a,b, 1998, 2000; Kirpotin et al., 1997b, 1998, 2000). Pharmacokinetic studies in rats demonstrated minimal differences in circulation lifetimes between nontargeted sterically stabilized liposomes and anti-HER2 immunoliposomes even after repeated weekly injections of immunoliposomes (t_{1/2} ~16 h). This is important because antibody conjugation has been previously shown to significantly reduce the circulation lifetimes of liposomes when attached directly to the liposome surface (Debs et al., 1987) or to the termini of liposome-conjugated PEG chains (Mori et al., 1991; Allen et al., 1994, 1995b; Goren et al., 1996; Zalipsky et al., 1996), especially after repeated administration (Harding et al., 1997). Evidently, the use of Fab' or scFv fragments for targeting eliminated unwanted interactions of extraneous IgG sequences with immune cells contributed to uncompromised pharmacokinetics of these immunoliposomes

of Gabizon and co-workers, who developed a similar immunoliposome somes (Kirpotin et al., 1997b, 2000). This was similar to the findings grafts to a greater extent than nontargeted sterically stabilized liposomes did not accumulate in HER2-overexpressing human tumor xenousing a noninternalizable antibody for tumor targeting (Goren et al., compared to nontargeted SSL and was often found within the canever, the distribution of liposomes within the tumor was more uniform accumulation of liposomes in the tumor and not active targeting. Howtumor and low rate of subsequent removal from the tumor control the cer cells themselves (Kirpotin et al., 1997b, 2000), similar to that seen 1996). These results suggest that the rates of extravasation into the Fujimori et al., 1990). to receptors located at the surface of the tumor (Weinstein et al., 1987. overcome the binding site barrier resulting from tight antibody binding immune fragments compared to full antibodies. This may help partially form distribution may be a result in part of the relatively low avidity of tumor-resident macrophages. The greater penetration and more unitargeted sterically stabilized liposomes were most often found within in HER2-overexpressing cells in culture (Kirpotin et al., 1997a). Non-An interesting finding was that anti-HER2-targeted immunolipo-

The favorable pharmacokinetics, tumor distribution, and endocytosis of anti-HER2 Fab' and scFv-immunoliposomes helped translate into higher antitumor efficacy of anti-HER2 liposomal doxorubicin in xenograft models of HER2-overexpressing human breast cancers in immunodeficient mice (Park et al., 1997a, 2000). Injections of HER2-targeted or nontargeted liposomal doxorubicin or free doxorubicin were started when tumor volumes reached 200–1000 mm³ and continued for total

V = A

were less effective than doxorubicin formulated into anti-HER2 im saline control, free doxorubicin, or nontargeted SSL in reducing tuof three weekly doses of 5 mg/kg. Antitumor efficacy was monitored considerable number of tumors, up to 60%, completely regressed with mor size following this schedule of treatments (Table III). In addition a Anti-HER2 immunoliposomes were shown to be more effective than at sizes equal to or less than 100 mm³ due to the poor responsiveness antitumor efficacy studies on xenografts, where treatment often starts in volume (Table III) (Park et al., 2000). This is an unusual result for Significant reduction in tumor size was achieved when the immunono pathological evidence of tumor cells in the tumor inoculation site by measuring the decrease in tumor size following start of treatment nontargeted liposomal doxorubicin. However, both these treatments the clinic against HER2-overexpressing breast cancers (trastuzumab overexpressing tumors (Baselga et al., 1999) and, in fact, is now used in cal to these liposome-conjugated Fab' suppresses the growth of HER2 munoliposomes. Because the immunoglobulin with Fab portions identitoxicities or weight loss in the nude mice treated with anti-HER2 imof larger tumors to most chemotherapy. There were also no notable liposome treatment was delayed until the tumor reached 1000 mm³ orubicin delivered in HER2-targeted immunoliposomes against HER2 munoliposomes (Park et al., 2000). Superior antitumor activity of dox HER2 immunoliposomes and the combination of trastuzumab IgG with Herceptin), control experiments were performed with "empty" antioverexpressing xenografts appears to be the result of their specific the criteria for ligandoliposome design laid out in Table II) 1998, 2000) and the overall in vivo performance of anti-HER2 immunointernalization by malignant cells in the tumor (Kirpotin et al., 1997b liposomes, as the targeted drug carrier was related to the fulfillment of

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E. TARGETED DELIVERY OF NUCLEIC ACIDS TO CELLS THROUGH HER2 RECEPTORS

Targeting gene delivery to malignant cells has also been shown to increase transfection efficiency using both viral (Goldman et al., 1997; Boerger et al., 1999; Gu et al., 1999) and cationic lipid-based (Kao et al., 1996; Kikuchi et al., 1996; Park et al., 1997b,c; de Lima et al., 1999) gene-delivery approaches. Although cationic lipid-based gene delivery vehicles can be quite efficient in vitro, the need to modify the lipid compositions for increased stability in vivo also reduces their transfection competency. Modifying the complexes for specific targeting to growth factor receptors has been shown to help overcome some of this loss in

ANTITUMOR ACTIVITY OF VARIOUS DOXORUBICIN FORMULATIONS AGAINST BT-474
XENOGRAFIS IN NUDE MICE^{a,b}

					Treatmer	Treatment outcome
		Treatme	Treatment started			Complete
		Tumor size,		Tumor size,		regressions/
Treatment	At day	(mm ³)	At day	(mm³)	B	total tumors
Study A						
Saline control	7	206 ± 25	43	4150 ± 2724		0/11
L _b -DOX	7	235 ± 38	4 3	425 ± 370		0/11
Anti-HER2	7	232 ± 24	43	105 ± 132	0.013 (vs	5/10
IL ₉ -DOX					Ls-DOX)	
(Fab')						
Anti-HER2	7	203 ± 32	43	50±33	0.006 (vs	6/11
ILs-DOX					La-DOX)	
(acfv)						
Study B						<u>:</u>
DOX +	25	999 ± 387	55	5100 ± 4035		0/10
Trastuzumab						
Anti-HER2	25	1031 ± 371	56	263 ± 175	0.006	0/10
ILs DOX						
Study C						:
Saline control	16	370 ± 58	£	3800 ± 1270		0/10
La-DOX+	16	353 ± 29	£	783 ± 344		0/10
Trastuzumab						
Anti-HER2	16	338 ± 29	43	295 ± 147	0.003	3/10
IL ₈ DOX						

^aAdapted from Park et al. (2000).

efficiency (Park et al., 1997b,c). Condensed and PEG-stabilized cationic lipid-DNA complexes were shown to have a 20-fold increase in reporter gene expression following addition of anti-HER2 Fab'-PEG-DSPE conjugates to the formulation (Kirpotin et al., 1998). Receptor-specific targeting has also been shown to increase the amount of complexed agent reaching the nucleus. Thus, using cationic lipid complexes of antisense oligonucleotides, a significant increase in nuclear localization of fluorescein-labeled oligonucleotides was seen when anti-HER2 Fab'-PEG-DSPE conjugates were added to the composition (Meyer et al., 1998).

The focus of this chapter has been on vitamin and growth factor receptors with respect to using them as targets for increasing binding and internalization of liposomes by malignant cells. However, liposomes and cationic lipid complexes can also be used to delivery therapeutic agents, such as antisense oligonucleotides or genes that code for inhibitory proteins or inhibit growth factor receptor expression by binding to nascent RNA (Wang et al., 1995; Hung et al., 1998; Muller et al., 1998). For example, both cationic lipid-DNA complexes and adenoviral vectors have been used to deliver the E1A gene and the nontransforming simian virus 40 large-T-antigen mutant gene to tumors in both mice and humans (Yu et al., 1995; Chang et al., 1997; Hortobagyi et al., 1998; Hung et al., 1998). Both genes are thought to code for gene products that can repress HER-2/neu promoter function.

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F. LIPOSOME TARGETING USING OTHER GROWTH FACTOR RECEPTORS

creased in vitro uptake of anti-EGFR IgG-immunoliposomes by EGFR with PEG, a portion of which was bearing terminal hydrazide groups. In recombinant anti-EGFR immunoglobulin C225 to the liposomes coated has been reported. Harding et al. (1997) conjugated periodate-oxidized expressing prostate cancer cells (DU-145) was demonstrated in vitro. et al., 1996). In both studies, the growth factor-targeted uposomes were terior of the liposome (Rosenberg et al., 1987; Ishii et al., 1987; Kikuchi also by using the hormone itself as a targeting ligand attached to the ex however, at higher IgG conjugation, the circulation half-life of the li-(100 nm) did not significantly reduce liposome circulation longevity: IgG conjugation at approximately 13 IgG/molecules per liposome shown to be efficiently endocytosed by the target cells in vitro, either The targeting of both the NGF receptor and EGFR were accomplished injections. No therapy studies or tumor-uptake data were presented highly immunogenic in rats, resulting in fast clearance upon repeated posomes decreased threefold. Anti-EGFR IgG-immunoliposomes were Liposome targeting to cells expressing NGFR, EGFR, and VEGFR

bAnimals were treated according to the following protocols (tumor inoculation at day 0). Study A: 0.125 mg of doxorubicin encapsulated in nontargeted sterically stabilized liposomes (Ls-DOX) or in anti-HER2 immunoliposomes (anti-HER2 Ils-DOX) injected intravenously at days 7, 14, and 21. Study B: 0.05 mg of doxorubicin solution (DOX) (maximum tolerated dose) or 0.125 mg of anti-HER2 ILs-DOX injected intravenously at days 25, 32, and 39; DXR group also received 75 μg of anti-HER2 antibody (Trastuzumab) intraperitoneally at days 25, 28, 32, 35, 39, and 42. Study C: 0.1 mg of doxorubicin as SL-DOX or as anti-HER2 Ils-DOX injected intravenously at days 16, 22, and 29; Ls-DOX group also received 7.5 μg of Trastuzumab antibody intraperitoneally at days 16, 19, 22, 25, 29, and 32. Saline controls: animals received injections of excipient (HEPES- or phosphate-buffered physiological saline) instead of the drug injections. Tumor size data are mean ±standard deviation; p, probability of the null hypothesis for the difference of the mean tumor volumes between drug treatment groups at the indicated day (by the independent t test). The number of complete regressions is calculated at day 56.

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et al., 1997). Adenovirus was genetically engineered to express FGF seins to FGFR-overexpressing cancer cells (Beitz et al., 1992; Goldman attempted in these studies, leaving a considerable amount of work to berg et al., 1987). However, no in vivo therapeutic approaches were quences at the cell-seeking receptor sites of the capside, thus redirecting be done both in vitro and in vivo. FGF has been used to target toxgood targeting ligand also for liposomes. Cells expressing VEGF recepmor drug delivery (70–120 nm), these data suggest that FGF would be a size of adenovirus (90 nm) is in the size range of liposomes used for tuviral tropism to FGF-overexpressing cells (Gu et al., 1999). Because the fibroblasts (Ishii $et\ al.,\ 1987)$ or PC12 pheochromocytoma cells (Rosenand Szostak, 1990). Although promising results were obtained in in-(Willis et al., 1998) developed by the process called SELEX (Ellington tor were also targeted using an unusual ligand, nucleic acid aptamers sulted in a dramatic decrease in circulation lifetimes when compared hibiting cell proliferation in culture, the addition of the aptamers resingle injections. However, the potential immunogenecity of such a systo nontargeted liposomes. A reduction in the number of aptamers attached to the liposome surface may increase circulation lifetimes for geted liposome delivery, but also emphasize the importance of overall liposome targeting instead of more traditional peptide/protein ligands. issues are resolved, aptamers may become a promising alternative for tem may limit its usefulness if multiple injections are required. If these ligandoliposome design to realize the advantage of targeting in practice. These studies demonstrate potential of growth factor receptors in tar-

As it is the case with vitamin receptors (Section II,G), a largely unexplored area lies in using hormone receptors for liposome targeting Pepplide hormones and hormone-releasing factors, including plasma-stable peptide analogs and peptidomimetics, may be used as ligands to deliver liposomal drugs to endocrine tissues and hormone-responsive nonendocrine tissues such as breast and uterus. Thus, a recent study successfully used doxorubicin conjugate with luteinizing hormone-releasing hormone (LH-RH) to deliver the cytotoxic drug to a human prostate cancer xenograft (Koppán et al., 1999). The experience of using these and other ligands reactive with hormone and growth factor receptors together with the principles of rational ligandoliposome design (Table II) will help to create a new generation of liposomal drug-delivery systems.

IV. CONCLUSIONS AND FUTURE PERSPECTIVES

Due to their unique place in providing for cellular growth and differentiation, and for their ability to internalize into the cell, vitamin

and growth factor receptors undergo changes in malignant cells that more liposome formulations entering the oncopharmaceutical market, enough to provide for a practically feasible ligandoliposome. Now, with bly because the field of liposome pharmacology has not been advanced in vitro studies in this area, little has been accomplished in vivo, possimake these cellular markers good recognition tags for targeted oncological drug delivery using liposomes. To date, despite many promising anchors) will allow these compounds to enter the clinic. In this chapthe development of new advanced technologies for making targeting ligands (scFv, aptamers) and their conjugates to liposomes (PEG-DSPE these exemplary cases are easy to extrapolate onto a vast variety of HER2 antibody fragment. The successful methodologies developed in liposome targeting systems directed to folate receptor via its natural illustrated how these principles were implemented in two successful ter the principles of ligandoliposome targeting were outlined; we also method in drug delivery. the time when ligand-directed liposome targeting becomes a routine ligands and lipid-based drug-delivery constructs, bringing us closer to ligand, folic acid, and to HER2/neu growth factor receptor via an anti-

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Vitamins and Homocysteine Metabolism

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Reference

I. INTRODUCTION

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Homocysteine is a thiol-containing intermediary metabolite that lies at a crossroads of three important metabolic processes: the methionine cycle, the folate cycle, and the transsulfuration pathway. In recent years, there has been increased interest in homocysteine metabolism because of the clinical observation that individuals with elevated levels of total plasma homocysteine (tHcy) are at increased risk of vascular disease and that woman with elevated tHcy levels are at increased risk of giving birth to children with neural tube defects.

Three vitamins play a key role in homocysteine metabolism: pyridoxine (B_6) , cobalamin (B_{12}) , and folate. Deficiency in any of these three vitamins can cause increased tHcy levels and increased risk of vascular disease and birth defects. In this chapter the current state of knowledge of homocysteine metabolism is summarized. The first section of this chapter discusses the molecular details of the role of these vitamins in the enzyme chemistry of homocysteine metabolism. The second section discusses clinical and epidemiologic data concerning the